# **FULL PAPER**

# **Thermal Ring-Opening Polymerization of Hydrocarbon-Bridged [2]Ferrocenophanes** : **Synthesis and Properties of Poly(ferrocenylethy1ene)s and Their Charge-Transfer Polymer Salts with Tetracyanoethylene**

# **James M. Nelson, Paul Nguyen, Ruth Petersen, Heiko Rengel, Peter M. Macdonald, Alan J. Lough, Ian Manners;\* Nandyala P. Raju, John E. Greedan, Stephen Barlow and Dermot O'Hare\***

Abstract: The poly(ferrocenylethylene)s  $[Fe(\eta-C_{5}H_{3}RCH_{2})_{2}]_{n}$  5a and 5b (a:  $R = H$ , **b**:  $R = Me$ ) have been prepared by thermal ring-opening polymerization of the corresponding strained hydrocarbon-bridged [2]ferrocenophanes  $[Fe(\eta-C_s-$ H,RCH,),] **(4a** and **4b). An** X-ray diffraction study of 4a indicated significant strain. Polymer **5a** was crystalline and insoluble in common organic solvents and was characterized by solid-state  ${}^{13}C$ NMR. Polymer **5b,** which was soluble in organic solvents, was characterized by 'H and **I3C** NMR, UV/visible spectroscopy and elemental analysis. Its molecular weight distribution was bimodal (gel permeation chromatography:  $M_w = 9.6 \times$ 

 $10^4$ ,  $M<sub>n</sub> = 8.6 \times 10^4$  for the high molecular weight fraction,  $M_w = 4.8 \times 10^3$ ,  $M<sub>n</sub> = 3.5 \times 10<sup>3</sup>$  for the oligomeric fraction), suggesting two polymerization mechanisms. The UV/visible spectrum implied a localized structure for the polymer backbone. Cyclic voltammetry revealed that **5 b** undergoes two reversible oxidations in CH<sub>2</sub>Cl<sub>2</sub> solution at  $-0.25$ and  $-0.16$  V. The redox coupling is indicative of only a small degree of interaction between the iron centres. Ther-

**Keywords**  ceramics ferrocenes \* iron - magnetic properties \* polymers

mogravimetric analysis indicated that **5a**  and **5b** are thermally stable to ca. 300- 350 °C under  $N_2$ . At higher temperatures they yield ferromagnetic iron carbide ceramics **6a** and **6b** (ca. 50% and 32%, respectively, at  $600^{\circ}$ C) together with molecular depolymerization products. The reaction of **5 b** with tetracyanoethylene (TCNE) yielded insoluble and soluble oxidized products **11** and **12,** which differed in the degree of oligomerization of the  $TCNE<sub>x</sub><sup>y-</sup>$  counterions. These products were characterized by IR, elemental analysis, ESR spectroscopy, and magnctic susceptibility measurements. The last revealed the presence of significant antiferromagnetic interactions in **12.** 

# **Introduction**

Transition-metal-based molecular and oligomeric materials and their associated charge-transfer salts are of considerable current interest for their redox chemistry and solid-state properties.<sup> $[1, 2]$ </sup> Transition-metal-containing polymers are also attracting growing attention as a consequence of their advantageous processability, their physical and catalytic properties, and their potential use as ceramic precursors. $[3-10]$  However, until recently progress in this area has been seriously impeded by the lack of viable synthetic routes to high molecular weight examples of

[\*] I. Manners, J. M. Nelson, P. Nguyen, R. Petersen, H. Rengel, P. M. Macdonald, A. J. Lough Department of Chemistry, University of Toronto 80 St. George **St.,** Toronto M5S3H6, Ontario (Canada)  $Fast: Int. code + (416)978-6157$ N. P. Raju. J. E. Greedan Institute for Materials Research, McMastcr University

Hamilton LSS4M1, Ontario (Canada) D. O'Hare, S. Barlow Inorganic Chemistry Laboratory, University of Oxford South Parks Road. Oxford OX13OR (UK) Fax: Int. code +(1865)272690

these materials. With this in mind, we reported the discovery that strained [l]ferrocenophanes with a single silicon atom in the bridge, such as **I,** undergo thermal ring-opening polymerization (ROP) to yield high molecular weight poly(ferrocenylsi1anc)s **2**  (Scheme 1).<sup>[11]</sup> We have subsequently shown that the corresponding germanium-, $^{[12]}$  phosphorus-, $^{[13]}$  sulfur-, $^{[14]}$  and tinbridged<sup>[15]</sup> [1]ferrocenophancs also polymerize thermally to yield high molecular weight poly(ferrocene)s.<sup>[16]</sup> Recent research has focussed on detailed studies of the synthesis and properties of these interesting materials. $[17 - 31]$ 

Polymerizable [l]ferrocenophancs possess strained structures in which the planes of the cyclopentadienyl (Cp) ligands are tilted by ca.  $14-31^\circ$  relative to one another.<sup>[13-15.32-37]</sup> How-



Scheme 1. Thermal ring-opening polymerization of a strained [1] ferrocenophane 1 with a single bridging silicon atom to yield poly(ferrocenylsilane) 2.

ever, our attempts to extend the ROP methodology to the disilane-bridged [2]ferrocenophane **3** were unsuccessful.[381 This was explained by the lower degree of ring strain present in this species, which is reflected by the very small tilt angle of  $4.19(2)$ <sup> $\degree$ </sup>.<sup>[38, 39]</sup> As part of our strategy to increase the polymerizability of [2]ferrocenophane systems, we studied the ROP chemistry of [2]ferrocenophanes with a hydrocarbon bridge **4a,b.** These species were prepared by the reaction of the dilithium reagents  $Li_2[C_5H_3RCH_2]_2$  (R = H, Me) with FeCl<sub>2</sub> in THF. As illustrated by the previously determined structure of 4c,



which possesses a Cp ring tilt angle of ca.  $23^{\circ}$ ,<sup>[40]</sup> these species are significantly more strained than **3** because of the smaller size of carbon relative to silicon. We subsequently reported that these hydrocarbon-bridged [2]ferrocenophanes undergo therinally induced ROP to yield poly(ferroceny1ethylcnc)s **5 a** and **5b** (Scheme 2).<sup>[41, 42]</sup> We found that the analogous [2]ruthenocenophanes also polymerize when heated.<sup>[43]</sup>





In this paper, we report full details of our work on the ROP of hydrocarbon-bridged [2]ferrocenophanes, and the characterization and properties of the resulting polymers. We also describe studies of the charge-transfer salts derived from the oxidation of **poly(ferrocenylethy1ene)s.** Although the electrochemistry of polymers with ferrocene units in their main chain has been the subject of several studies, few recent investigations have been madc involving *chrmical* oxidation of such polymers. Brandt and Rauchfuss have reported briefly that bromine oxidizes carbon disulfide solutions of high molecular weight poly(ferrocenylene persulfide)  $[Fe(\eta - C_5H_3nBu)(\eta - C_5H_4)S_2]$  to yield a soluble black material of approximate formula  $[Fe(\eta - C_5H_3$  $nBu$ )( $\eta$ -C<sub>5</sub>H<sub>4</sub>)S<sub>2</sub>]<sub>n</sub>Br<sub>0.5n</sub>.<sup>[7a]</sup> Pannell, Diaz and coworkers have used UV/vis spectroscopy to study the oxidation of high molecular weight **poly(ferrocenyldialky1silane)s** by iron(m) chloride.<sup>[26]</sup> We have reported similar studies with  $FeCl<sub>3</sub>$  and, in addition, iodine and o-quinone oxidants and have shown that I,-oxidized samples of the high polymer **2** are localized on the  $57$ Fe Mössbauer timescale at room temperature.<sup>[44]</sup> In a recent, particularly intriguing study by Garnier et al., the oxidation reactions of the low molecular weight oligo(ferroceny1 dialkylsilane)s with tetracyanoethylene (TCNE) in dichloromethane were reported to yield dark precipitates; in each case analysis indicated these precipitates contained one TCNE per monomer iron unit, and <sup>57</sup>Fe Mössbauer data showed they contained a mixture of Fe" and Fe"' sites, which were localized at 80 K and in rapid exchange at room temperature.<sup>[27]</sup> Although IR data assigned to neutral and ionic polymer units were reported, the  $C \equiv N$  stretches of the cyanocarbon counterions were not discussed. The authors interpreted the magnetic data as indicative of both ferromagnetic and spin glass behaviour. This is quite surprising given that one would expect the first oxidation potentials of these materials to be close to those of ferrocene and thc fact that ferrocene forms a complex with TCNE, which only dissociates to ionic species in polar solvents. The observation of room-temperature Mössbauer detrapped behaviour in this class of bridged metallocene system is also highly unusual.

We have recently described the oxidation reactions of a number of high molecular weight poly(ferrocenylsilane)s with  $TCNE<sub>1</sub><sup>{45}</sup>$  which were carried out in analogous fashion to those described by Garnier et al.<sup>[27]</sup> Poly(ferrocenylsilane)s lacking ring methylation gave no reaction with TCNE; this contrasts with the results reported by Garnier et al. for apparently analogous low molecular weight oligomers. However, cyclopentadienyl methylated polymers did react with TCNE, consistent with the effect of methylation upon their oxidation potentials. IR. spectroscopy showed unusual cyanocarbon anions to be present. In this paper we also describe the analogous reaction between poly(ferrocenylethylene)s and TCNE.

# **Experimental Section**

**Materials:** Dicyclopentadiene. methylcyclopcntadicne dimer. hexamethylphosphoramide (HMPA) and 1.2-dibromoethane were purchased from Aldrich and were distilled before use. Iron( $\mathfrak n$ ) chloride (Aldrich), sodium metal (Aldrich), 1.6<sub>M</sub> nBuLi in hexanes (Aldrich), PtCl, (Strem) and platinum divinyltetramethyldisiloxane complex in xylene (United Chemical Technologics) were used as received. Bis(cyclooctene)rhodium(i) chloride dimer<sup>[46]</sup> and the dilithium salts  $Li_2[MeC_5H_3CH_2]_2$  and  $Li_2[C_5H_4CH_2]_2^{(47)}$ wcrc prcparcd by literature methods. TCNE (Aldrich) was purified by vacuum sublimation.

**Equipment:** All rcactions and manipulations were carried *out* under an *atmo*sphcrc of purified nitrogen either by means of Schlenk techniques or in an inert-atmosphcrc glovcbox (Vacuum Atmospheres). except for the purification of the polymers 5a and 5b, which was carried out in air. Solvents were dricd by standard methods. distilled, and then stored under nitrogen over activated molecular sieves. The 200 or 400 MHz <sup>1</sup>H NMR spectra and 50.3 or 100.5 MHz *"C* NMR spectra were rccordcd on a Varian Gemini 200 or Unity 400. NMR chemical shifts were referenced to residual protonated solvent peaks. Solid-state <sup>13</sup>C NMR spectra were obtained with a Chemagnetics CMX 300 spectrometer equipped with a Chemagnetics magic angle spinning probe doubly tuned to the resonance frequencies of 13C *(75.3* MHz). Samples were spun in a 7.5 mm o.d. zirconium rotor at a spinning rate of 60110 Hz. **A** single-contact cross-polarization technique was employed with a contact time of *5* **ms** and proton decoupling during the signal acquisition. The proton radial frequency field strength was 50 kHz. Spectra were acquired with a sweep width of 50 kHz, a data size of 2 Hz and a recycle delay of 5 s. All chemical shifts were referenced to external TMS. Room-temperature <sup>57</sup>Fe Mössbauer spectra were obtained by means of a Ranger Scientific Inc. VT-1200 instrument with a MS-1200 digital channel analyzer. The *y* source was a 6 mCi <sup>57</sup>Co sample supplied by Amersham. The data were collected in  $a - 15.8$  mm s<sup>-1</sup> to  $+15.8$  mm s<sup>-1</sup> range and referenced to Fe foil and processed as described previously. Mass spectra were obtained with the use of a VG 70-250s mass spectromctcr operating in Electron Impact (EI) mode. Molccular weights were estimated by gel permeation chromatography (GPC) on a Waters Associates liquid chromatograph equipped with a 510 HPLC pump, U6K injector, ultrastyragel columns with a pore size between  $10^3$ -10<sup>5</sup> Å, and a Waters 410 differential refractometer. A flow rate of 1.0 mL min<sup>-1</sup> was used and samples were dissolved in a solution of 0.1% tetra-n-butylammonium bromide in THE Ten samples of monodisperse polystyrene in the range  $M_w = 10^3 - 10^6$  were used as standards for calibration purposes. Elemental analyses were performed by Canadian Microanalytical Service, Delta, B. C (Canada) and Quantitative Technologies, Whitehousc, NJ (USA) or by the Analytical Department of the Inorganic Chemistry Laboratory, Oxford (UK). UV/Visible spectra were recorded on a Hewlett-Packard 8452 A Diode Array Spectrophotometer with a 1 cm quartz cell: the  $\epsilon$  values quoted have the units  $L \text{mol}^{-1} \text{cm}^{-1}$  and for the polymers are per repeat unit. IR spectra were recorded on a Mattson Intruments Polaris spectrometer.

A Perkin Elmer DSC-7 differential scanning calorimeter equipped with a TAC7 instrument controller was used to study thermal bchaviour. The thermograms were calibrated with the melting transitions of decane and indium and were obtained at a heating rate of 10 °Cmin<sup>-1</sup> under dinitrogen. A Perkin- Elmer TGA-7 thcrmal gravimetric analyzer equipped with a TAC 7 instrument controller was used to study polymer thermal stability. The thermograms were calibrated with the magnetic transitions of Nicoseal and Perkalloy and were obtained at a heating rate of  $10$   $°C$ min<sup>-1</sup> under dinitrogen .

Electrochemical experiments were carried out on a PAR model 273 potcntiostat with a Pt working clcctrode, a W secondary electrode, and an Ag wire reference electrode in a Luggin capillary. Polymer solutions were  $5 \times 10^{-3}$  M in CH<sub>2</sub>Cl<sub>2</sub> with 0.1 M [Bu<sub>4</sub>N][PF<sub>6</sub>] as supporting electrolyte. Peak currents were found to be proportional to the square root of the scan rate over the range studied (25 to 1000 mVs<sup>-1</sup>); this indicates that charge transfer is similar to a semi-infinite linear diffusion process.

Wide-angle X-ray scattering data wcrc obtained with a Siemens D 5000 diffractomer employing Ni-filtered Cu<sub>Kz</sub> ( $\lambda = 1.54178$  Å) radiation. The samples were scanned at step widths of  $0.02^{\circ}$  with 1.0 s per step in the Bragg angle range of  $5-90^\circ$ . Samples for the X-ray studies were prepared by spreading the finely ground polymer on grooved glass slides.

High-resolution SEM with EDX and BE1 were carried out by Imagetak Analytical Imaging. Errors in thc compositional values obtained are considered to be  $\pm 5\%$  for Fe and  $\pm 10\%$  for C. XPS data was collected on a Leybold MAX200 instrument. Values obtaincd are based on the integration of the peaks Fc (2p, *312,* 708.1 eV) and *C* (1 **s,** 284.6 eV).

**All** pyrolyses were carried out under an atmosphere of prepurificd nitrogen in a 36" 3-zone Lindberg Pyrolysis Oven (Model 55035) with a  $1<sup>5</sup>/<sub>8</sub>$ " internal diameter and Thermcraft control system Model 3D1-50-115 (UP27) with type K thermocouples and independent temperature control. **A** program was created that ramped to the dcsircd temperature over a **1** h period. Polymer samples were loaded in quartz boats,  $2 \times \frac{1}{2} \times \frac{1}{2}$ , and inserted into quartz pyrolysis tubes 36' long with a 1" external diameter, equipped with quartz liners 32" long with a 1" external diameter. Magnetization measurements were performed with a Quantum Design SQUID magnetometer.

ESR measurements were performed in high purity Spectrosil<sup>TM</sup> quartz tubes with an X-band Varian spectrometer; peaks were referenced by means of a microcrystalline sample of *1,1* -diphenyl-2-picrylhydrazyl. Solid-state magnetic susceptibilities were measured on samplcs loaded in gelatin capsules with a Quantum Design MPMS-7 SQUID magnetometer, operated at fields of 0.1 T.

**Synthesis of the** *[2]ferrocenophane* 4a: A solution of  $Li_2[C_5H_4CH_2]$ , (0.35 g, 2.07 mmol) in THF (40 mL) was added dropwise 10 a suspension of FcCI, (0.26 g, 2.07 mmol) in the same solvent (40 mL) at  $-78$  °C. The reaction mixture was stirred at this temperature for 3 h and was then allowed to warm to room temperature over a 12 h period. Following solvent removal in vacuo, dark red microcrystalline **4a** was isolated and purified by vacuum sublimation (80"C, 10mmHg). Yield 0.285g (65%): m.p. 120 'C; 'HNMR (200 MHz)  $(C_6D_6)$ :  $\delta = 4.6$  (t, 4H, Cp), 3.9 (t, 4H, Cp), 2.6 (s, 4H, CH<sub>2</sub>CH<sub>2</sub>); <sup>13</sup>C NMR (C<sub>6</sub>D<sub>6</sub>):  $\delta$  = 91.2 (ipso, Cp), 68.8, 72.9 (x and  $\beta$ Cp), CH<sub>2</sub>CH<sub>2</sub>); <sup>13</sup>C NMR (C<sub>6</sub>D<sub>6</sub>):  $\delta$  = 91.2 (ipso, Cp), 68.8, 72.9 (x and  $\beta$ Cp), 34.1 (CH<sub>2</sub>CH<sub>2</sub>); UV/Vis (THF):  $\lambda_1 = 474$  ( $\vare$ 34.1 (CH<sub>2</sub>CH<sub>2</sub>); UV/Vis (THF):  $\lambda_1 = 474$  ( $\varepsilon_1 = 450$ ),  $\lambda_2 = 268$  (sh,  $\varepsilon_2 = 2250$ ),  $\lambda_3 = 220$  nm  $(\varepsilon_3 = 18500 \text{ m}^{-1} \text{ cm}^{-1})$ ; MS (E1, 70 eV):  $m/e = 212$  $(M^+, 100\%)$ ; elemental analysis for C<sub>12</sub>H<sub>12</sub>Fe: calcd. C 67.9, H 5.6%; found C 67.9, H 5.7%.

**Synthesis of the [2] ferrocenophane 4b: A solution of**  $Li_2[C_5H_3(CH_3)CH_2]$  $(0.50 \text{ g}, 2.52 \text{ mmol})$  in THF  $(40 \text{ mL})$  was added dropwise to a suspension of FeCl<sub>2</sub> (0.32 g, 2.52 mmol) in the same solvent (40 mL) at  $-78$  °C. The reaction mixturc was stirred at this temperature for- 3 h and was then allowed to warm to room temperature over a 12 h period. Following solvent removal in vacuo, **4b** was isolated as a viscous red oil and purified by vacuum distillation (120 °C, 10 mm Hg). Yield 0.45 g (74%). MS (EI, 70 eV):  $m/e = 240$  ( $M^+$ , 100%), 225 *(M<sup>+</sup>* – CH<sub>3</sub>, 45%); <sup>1</sup>H NMR (200 MHz, C<sub>6</sub>D<sub>6</sub>):  $\delta = 4.4-4.7$ (brm, 3H, Cp), 3.7-4.1 (brm, 3H, Cp), 2.5-2.7 (s, 4H, CH<sub>2</sub>CH<sub>2</sub>), 1.7-2.2 (s, 6H, Me); <sup>13</sup>C NMR (100.5 MHz, C<sub>6</sub>D<sub>6</sub>):  $\delta = 85.6 \cdot 91.5$  *(ipso, Cp), 67.0* 80.7 ( $\alpha$ ,  $\beta$ Cp), 32.3-34.8 (CH<sub>2</sub>CH<sub>2</sub>), 13.9-15.5 (Me). Because of the existence of different isomers the NMR spectra of **4 b** consisted of numerous *(2* 7) peaks in the regions indicated. UV/Vis (THF):  $\lambda_1 = 470$  ( $\varepsilon_1 = 456$ ),  $\lambda_2 = 268$ (sh,  $\varepsilon_2=2350$ ),  $\lambda_3=218$  nm ( $\varepsilon_3=18900$  M<sup>-1</sup> cm<sup>-1</sup>).

**Ring-opening polymerization of 4a and 4b; synthesis of the poly(ferrocenylethylene)s 5a and 5b:** Polymers 5a and 5b were prepared similarly and the general synthesis **ia** illustrated by thal of **5b.** 

A sample of 4b (1.00 g, 4.2 mmol) was allowed to polymerize in an evacuated, sealed Pyrex tube at 300°C for 1 h. The tube contents became molten and rapidly more viscous and, after 1 h, complctcly immobile. The polymeric product was dissolved in THF (40 mL) over 1 h and the resulting solution concentrated to *5* mL. This was then added dropwise to a large excess of methanol to yield **5b** as a mustard-coloured solid. Yield 0.95 g (95%). After multiple precipitations from THF into methanol, the yields of purified **Sb**  were generally in the range of 80 -95%. Polymerization times varied but were usually  $\approx$  45 min-1 h. In some cases the polymer product was yellow-brown. particularly in cases where heating lasted for > 1 h. This is probably *a* consequence **of** small amounts of thermal dccomposition (scc polymer pyrolysis experiments below).

**5a:** Yellow-brown insoluble material, which was obtained in the form of film directly from the polymerization tube. Solid-state <sup>13</sup>C CP-MAS NMR:  $\delta = 90.3$  (ipso, Cp), 70.1 (x,  $\beta$ Cp), and 37.0 (CH<sub>2</sub>CH<sub>2</sub>). Polymer 5a was extracted with hot  $CH_2Cl_2$ . Mass spectrometric analysis of the soluble material from the light orange solution identified the cyclic oligomcrs  $[Fe(C_5H_4CH_2)_2]_x$  (x = 2 - 5): MS (EI, 70 eV):  $m/e = 1060$  (x = 5, 18%), 848 Soxhlet extraction of polymer 5a in THF for 72 h was successful in producing a small amount of soluble material as a mustard-yellow fibrous powder: <sup>1</sup>HNMR (200 MHz, C<sub>6</sub>D<sub>6</sub>):  $\delta = 3.6-4.1$  (br, 8H, Cp), 2.3-2.8 (br, 4H, CH<sub>2</sub>CH<sub>2</sub>); GPC for THF-soluble extract: for first fraction,  $M_w = 8.1 \times 10^4$ ,  $M<sub>n</sub> = 6.6 \times 10<sup>4</sup>$ , polydispersity = 1.2. For second fraction,  $M<sub>w</sub> = 4.8 \times 10<sup>3</sup>$ ,  $M<sub>n</sub> = 3.5 \times 10<sup>3</sup>$ , polydispersity = 1.4. **(X** = *4,* **1S%!),** 636 *(X* = 3, 32%). 424 **(X** = *2,* 100°/u), 212 *(5* = 1. 90%).

**5b:** Mustard-yellow to yellow-brown fibrous powder 'H NMR (200 MHz,  $C_6D_6$ ):  $\delta = 3.6 - 4.1$  (br, 6H, Cp), 2.3 - 2.8 (br, 4H, CH<sub>2</sub>CH<sub>2</sub>), 1.6 - 2.1 (br, 6H, Me); <sup>13</sup>C NMR (100.5 MHz, C<sub>6</sub>D<sub>6</sub>):  $\delta = 83.3-88.5$  *(ipso, Cp), 67.0* 72.5 ( $\alpha$ ,  $\beta$ Cp), 31.4-33.4 (CH<sub>2</sub>CH<sub>2</sub>), 13.9-15.5 (Me). Because of the existence of different isomers, the I3C NMR spectra of **Sh** consisted of numerous  $(\geq 7)$  peaks in the regions indicated. The resonances for the different isomers were unresolved in the <sup>1</sup>HNMR spectrum. GPC: for first fraction,  $M_w =$  $9.6 \times 10^4$ ,  $M_p = 8.6 \times 10^4$ , polydispersity = 1.1. For second fraction,  $M_w =$ 4.8 × 10<sup>3</sup>,  $M_n = 3.5 \times 10^3$ , polydispersity = 1.4. UV/Vis (THF):  $\lambda_1 = 440$  $(\varepsilon_1 = 190)$ ,  $\lambda_2 = 268$  (sh,  $\varepsilon_2 = 3000$ ),  $\lambda_3 = 218$  nm  $(\varepsilon_3 = 20000 \text{ m}^{-1} \text{ cm}^{-1})$ ; clemental analysis for  $C_{14}H_{16}Fe$ : calcd. *C* 70.0, H 6.7; found *C* 69.8, H 6.8%.

### **Mechanistic investigations of the ring-opening polymerization of 4 b:**

 $(a)$  *Influence of polymerization time on the molecular weight distribution:* Sealed Pyrex tubes containing **4b**  $(\approx 0.2 \text{ g})$  were heated at 300 °C for i) 15 min, ii) 30 min, iii) 45 min, iv) **1** h, v) 1.5 h and vi) 2.5 h, at which time the contents of each tube werc examined by GPC in THF. i) The tube contents werc free-flowing and GPC analysis showed no signs of a substantial molecular weight fraction  $(M_w>1000)$ . ii) After 30 min the tube contents wcrc viscous and mobile and GPC analysis showed that the polymer present possessed an approximate weight-average molecular weight  $(M<sub>w</sub>)$  of  $8.0 \times 10<sup>4</sup>$ and a number-average molecular weight  $(M_n)$  of 6.6  $\times$  10<sup>4</sup> with no substantial lower molecular weight oligomeric fraction. iii) After 45 min the tube contents were immobile and GPC analysis showed a broad bimodal molecular weight distribution: for the high polymer fraction,  $M_w = 8.1 \times 10^4$ ,  $M_n = 6.3 \times 10^4$ , polydispersity = 1.3, and for the oligomeric fraction,  $M_w = 4.8 \times 10^3$ ,  $M_n = 3.5 \times 10^3$ , polydispersity = 1.4. iv) After 1 h the tube contents were immobile and GPC analysis showed a broad bimodal molecular weight distribution. GPC for the polymeric fraction,  $M_w = 7.8 \times 10^4$ ,  $M_n = 6.2 \times 10^4$ , polydispersity = 1.3; for the oligomeric fraction:  $M_w =$  $3.8 \times 10^3$ ,  $M<sub>n</sub> = 2.7 \times 10^3$ , polydispersity = 1.4. v) After 1.5 h the tube contents were immobile and slightly darkened in colour and GPC analysis showed a broad bimodal molecular weight distribution. GPC for the polymeric fraction:  $M_w = 7.8 \times 10^4$ ,  $M_n = 6.2 \times 10^4$ , polydispersity = 1.3; for the oligomeric fraction:  $M_w = 5.5 \times 10^3$ ,  $M_n = 3.5 \times 10^3$ , polydispersity = 1.6. vi) After 2.5 h the tube contents were immobile and contained a small insoluble fraction (ca. *5%).* GPC analysis of the soluble portion showed a broad bimodal molccular weight distribution. GPC for the polymeric fraction.  $M_w = 2.1 \times 10^4$ ,  $M_n = 1.6 \times 10^4$ , polydispersity = 1.3; for the oligomeric fraction,  $M_w = 1.7 \times 10^3$ ,  $M_n = 1.2 \times 10^3$ , polydispersity = 1.4.

*b*) *Influence of heating on the molecular weight distribution: In order to deter*mine whether the lower molecular weight fraction is produced as a result of thermal decomposition of the polymer. a tube containing a purified 0.1 g sample of polymer 5b was heated for 1 h at 300 °C. Analysis of the tube contents after this time by GPC showed no significant change in the molecular weight or in the molecular weight distribution

**Attempted transition-metal-catalyzed ROP of 4a:** To **a** solution **of4a** (20 mg, 0.09 mmol) was added bis(cyclooctene)rhodium(I) chloride dimer ( $\approx$  2 mg) in  $C_6D_6$ . This mixture was stirred under nitrogen for 48 h with constant monitoring of the reaction by <sup>1</sup>HNMR spectroscopy. Analysis of this mixture after 48 h showed that no reaction had occurred and no signs of insoluble **Sa**  were apparent. A 'H NMR spectrum of the mixture displayed characteristic resonances for the starting compound **4a.** <sup>1</sup>HNMR (200 MHz) (C<sub>6</sub>D<sub>6</sub>):  $\delta = 4.6$  (t, 4H, Cp), 3.9 (t, 4H, Cp), 2.6 (s, 4H, CH<sub>2</sub>CH<sub>2</sub>). Analysis of the reaction mixture by GPC showed no material of substantial molecular weight  $(M_{\rm w} > 1000)$ . Similar results were obtained with PtCl<sub>2</sub> and Pt divinyltetramethyldisiloxane catalysts.

#### **Pyrolysis of the poly(ferrocenylethy1ene)s Sa and** *S* **b under dinitrogen:**

*UJ TGA srtrdies:* A sample of **Sa** (0.02 g) was lightly packed into a TGA pan. which was then inserted in the TGA instrument under a steady flow of nitrogen for approximately one minute. The system was then heated from 25 to 600 °C at a heating rate of 10 °Cmin<sup>-1</sup>. When the program was complete and the furnace was allowed to cool to rooni temperature a black, lustrous ccrnrnic product **6a** was formed. yield 0.01 g (50%). This material was found to be readily attracted to a bar magnet. Similar studies with *5* **b** give a ceramic yield **of** 32%).

*h) Pyrolysis of 5b in a tube furnace:* A sample of 5b (0.50 g) was lightly packed into a quartz boat, which was then inserted into a pyrolysis tube. The tube was purged with a steady flow of nitrogen for approximately one minute. The system was then heated from 25 to 600 C over 1 h and *was* then maintained at a constant temperature of 600 °C for a further 4 h. During pyrolysis an orange-red liquid condensed on the cooler parts of the pyrolysis tube downstream from the quartz boat. When the program was complete and the furnace allowed to cool to room temperature, a black, lustrous ceramic product 6b was formed, yield 0.16 g ( $\approx$  30%). This material was also found to be readily attracted to a bar magnet. For ceramic **6b** (derived from *S* **b** at 600 C): for surface. XPS: Fc 1, *C* 87, Si 1. 0 11 %. For bulk. EDX: Fe 72, C 24, O 4%. Localized electron-rich sites. EDX: Fe 70, C 5, O 25%. Mössbauer spectroscopy: sextuplet IS = 0.03, MHS = 1.81 mm s<sup>-1</sup>. Magnetization measurements:  $H_c = 257$  G,  $M_r = 0.071 \mu_B$ /Fe,  $M_s = 0.17 \mu_B$ /Fe.  $XRD$ : sharp peaks at *d* spacing of 2.025, 1.429, and 1.167 Å assigned to  $\alpha$ -Fe. Other peaks together with several broadened peaks of low intensity (3.396, 2.956. 2.519, 2.467. 2.138. 2.300. 1.973: 1.850. 1.613, 1.482A).

The red-orange sublimate was collected by rinsing the pyrolysis tube with hexancs and was also analyzed. Mass spectrometry of the red-orange solution showed that it contained an inseparable mixture, which has been tentatively assigned to the linear methylated compounds  $[Fe(\eta-C_5H_3Me_2)(\eta-C_5H_3 (Me)(CH<sub>2</sub>)$ ], (MS (EI. 70 eV):  $m/e = 482$  ( $M<sup>+</sup>$ ,  $x = 2$ , 40%), 241 ( $x = 1$ , 100%)) and the cyclic trimer  $[Fe(\eta-C_5H_3(Me)CH_2)_2]_3$  (MS (EI, 70 eV):  $m/$ *<sup>P</sup>*=720 *(M* ', *S'%)).* 

**Reaction of the poly(ferrocenylethy1ene) Sb with TCNE:** Experiments were prrformed on samplcs of **Sb** that had been purified anaerobically. A solution of TCNE (138 mg, 1.08 mmol) in 10 mL CH<sub>2</sub>Cl<sub>2</sub> was added to a solution of purified polymer  $5b$  (250 mg, 1.04 mmol) in  $15 \text{ mL } CH$ , Cl<sub>2</sub>. The solution instantly darkened and a dark precipitate slowly appeared. After 12 h the solid was filtered off, washed with 50 mL  $CH<sub>2</sub>Cl<sub>2</sub>$  and dried in vacuo to yield a black powder, 11 (60 mg). Analysis (%): found (calcd. for the idealized composition [Fe(C<sub>5</sub>H<sub>3</sub>MeCH<sub>2</sub>)<sub>2</sub>]<sub>n</sub>[TCNE]<sub>n</sub>) *C* 66.41 (65.24), H 4.24 (4.38), N 13.20 (15.22), Fe 13.58 (15.17); ESR (solid 6.6 K): cation  $g_1 = 1.70$ .  $g_{\parallel} = 3.76$ ,  $\langle g \rangle = 2.58$ ; anion  $g = 2.00$ . Selected IR data (nujol mull): 2153 (br),  $2198$  (br) cm<sup>-1</sup>. The soluble portion was concentrated to 15 mL and added dropwise to vigorously stirred diethyl ether (175 mL). The precipitate was collected by filtration, washed with dicthyl ether (100 mL) and dried in vacuo to yield a green powder (95 mg), 12. Analysis (%): found (calcd. for the idealized composition  $[Fe(C_5H_3MeCH_2)_2]_n[TCNE]_n$ ) C 71.11 (65.24), H 4.86 (4.38). N 13.00 (15.22). Fe 12.55 (15.17); SQUID:  $\mu_{eff} = 1.7 \mu_B$ .  $\theta = -11.6 \text{ K}$ ; ESR (CH<sub>2</sub>Cl<sub>2</sub> glass, 6.8 K): cation  $g_1 = 1.70$ ,  $g_{\text{H}} = 3.81$ ,  $\langle g \rangle = 2.60$ ; anion  $g = 2.00$ ; selected 1R data (nujol mull): 2148 (br), 2199 (br), 2218 (br)  $cm^{-1}$ 

# **Results and Discussion**

Ferrocenyl groups have been introduced into the side-group structure of a wide range of organic macromolecules. $[48]$  Poly-(ferrocenylmethylene)s, of reported structure  $[Fe(\eta-C_5H_4),$ - $CH<sub>2</sub>$ , have been previously prepared by the zinc chloride/hydrogen chloride-catalyzed polymerization of ((dimethylamino) methyl)ferrocene reported by Neuse and Quo.<sup>[49]</sup> These materials are generally of low molecular weight  $(<10000)$  and have been found to exist as a mixture of  $1,2$ -,  $1,3$ -, and  $1,1'$ -disubstituted ferrocene units in the main chain.<sup>[48]</sup> To our knowledge, no well-characterized poly(ferroceny1ethylene)s of substantial molecular weight have bccn reported although oligomers as well as cyclics are formed in the synthesis of  $4a$ <sup>[50,51]</sup> In our initial communication,[411 we reported the thermal ROP of hydrocarbon-bridged [2]ferrocenophanes as a route to high molecular weight poly(ferrocenylethylene)s. In this paper, we discuss in detail the synthesis and properties of the resulting poly(ferrocenylethy1ene)s and thc characteristics of their charge-transfer salts with TCNE.

**Synthesis and characterization of the hydrocarbon-bridged [Z'jferrocenophanes 4 a and 4 b:** Ferrocenophanes containing hydrocarbon bridges have been known since the initial report of the methylated ethylene-bridged monomer **4c** by Burke Laing and Trueblood,<sup>[40]</sup> which was prepared by the coupling of 6,6dimethylfulvene with Na in THF to give the bisfcyclopentadieny1)tetramethylcthanc dianion, followed by the addition of FeCl, . The hydrocarbon-bridged [2]ferrocenophane **4 c** possesses a substantially ring-tilted structure, as indicated by singlecrystal X-ray diffraction studies, which showed that the cyclopentadienyl ligands are significantly tilted by  $\approx 23^\circ$  with respect to one another.

In the work dcscribed in this paper, compounds **4a** and **4b**  (which consists of a mixture of isomers) were preparcd by reaction of the dilithium reagent  $Li<sub>2</sub>[(C<sub>5</sub>H<sub>3</sub>RCH<sub>2</sub>)<sub>2</sub>]$  with iron(II) chloride in THF solutions (Scheme 3). Lentzner and Watts<sup>[52]</sup> initially reported this synthesis of **4 a;** however, these re-



Scheme 3. Preparation of compounds **4a** and **4b** by reaction of  $Li_2[(C_5H_3RCH_2)_2]$ with iron(ii) chloride in THF.

searchers experienced difficulty achieving high yields (yield = 3 *YO)* of the starting dilithium compound, owing to the formation of the spirocyclic by-product, **spiro[2,4]hepta-4,6-diene.**  The dilithium reagents used here were preparcd by the improved method reported by Collins and coworkers,<sup>[47]</sup> which involves the addition of hexamethylphosphoramide (HMPA) to the NaCp/1,2-dibromoethane mixture. The [2]ferrocenophanes were isolated as red, moisture-sensitive materials in  $\approx 70\%$  yield by vacuum sublimation or distillation. The structures of the [2]ferrocenophanes **4a** and **4b** were confirmed by <sup>1</sup>H and <sup>13</sup>C NMR spectroscopy and by mass spectrometry. This afforded data similar to that reported for the previously prepared methylated [2]ferrocenophane compound **4c.** In the case of **4a,** the molecular structure was also determined by single-crystal X-ray diffraction (see below). Thc UV/visibie spectra of **4 a** and **4 b** in the 200-800 nm range contained low-energy bands in the visiblc region at 470–474 nm ( $\varepsilon = 450 \,\mathrm{M}^{-1} \,\mathrm{cm}^{-1}$ ) that showed a characteristic bathochromic shift and increase in intensity relative to the long-wavelength band of ferrocene at  $440 \text{ nm}$  ( $\varepsilon =$  $90<sup>-1</sup>$  cm<sup>-1</sup>). Osborne and co-workers have previously commented that this is indicative of significant tilting of the cyclopentadienyl rings.[531 Similar bathochromic shifts were also detected for the methylated [2]ferrocenophane **4c** ( $\lambda = 472$  nm,  $\varepsilon = 400 \,\mathrm{m}^{-1} \,\mathrm{cm}^{-1}$ ), which was found to possess a significantly ring-tilted structure with a tilt angle of  $23(1)^{\circ}$ .<sup>[52]</sup>

**Discussion of the X-ray structure of 4a:** Several [2]ferrocenophanes have been crystallographically characterized, including the previously mentioned methylated [2]ferrocenophane  $4c^{[40]}$  and the unstrained disilane  $3$ ,<sup>[38,39]</sup> as well as the S-N bridged [2]thiazaferrocenophane **7** reported by Abramovitch



and co-workers,<sup>[54]</sup> which was found to possess a substantially strained, ring-<br>tilted structure (tilt angle of<br>23°). Very recently, Hafner and coworkers<sup>[55]</sup> published thc synthesis and Xray structural analysis of a doubly strapped hydrocar-

bon-bridged [2]ferrocenophane **8**  $(R = R' = H)$ , which was found to possess a considerably ring-tilted structure with a tilt angle of 28.8". In order to study further the influence of a hydrocarbon bridge on the strain and polymerizability of these ferrocenophane systcms, an X-ray crystallographic study of **4a** was carried out. Crystals of **4a** were isolated from hexanes solution at  $-20$  °C. The molecular structure of **4a** is shown in Figure 1. **A** summary of cell constants and data collection parameters are included in Table 1, and important bond lengths and angles are listed in Table 2. The angles  $\alpha$ ,  $\beta$  and  $\delta$  used in discussing the structures are defined in Figure *2.* The structure of **4a** is disordered over two sites  $(C(12a), C(12b))$  with occupancies of 0.60 and 0.40.

Table 1. Summary of crystal data and intensity collection parameters for 4a [a].



[a] Crystallographic data (excluding structure factors) for the structure reported in this paper have becn deposited with thc Cambridge Crystallographic Data Centre as supplementary publication no. CCDC-100 124. Copies of the data can be obtained free of charge on application to The Director, CCDC, 12 Union Road, Cambridge CB21EZ. UK (Fax:  $Int. \text{code} + (1223)336-033; \text{e-mail}$ : de-Cambridge CB21EZ, UK (Fax: Int. code + (1223) 356-035; e-mail: de-<br>posit@chemcrys.cam.ac.uk). [b]  $R_1 = \sum (F_o - F_c)/\sum (F_o)$ . [c]  $wR_2 = [\sum [w(F_o^2 F_c^2$ <sup>2</sup>]/ $\sum [w (F_o^2)^2]$ ]<sup>1/2</sup>; see ref. [71].

Table 2. Selected distances  $(A)$  and angles  $(\degree)$  for **4a** (major component).

| $Fe-C1$   | 1.965(6)  | $Fe-C6$  | 1.966(6)   |
|---|---|--|--|
| $Fe-C2$   | 2.022(6)  | $Fe-C7$  | 2.007(6)   |
| $Fe-C3$   | 2.060(5)  | $Fe-C8$  | 2.054(6)   |
| $Fe-C4$   | 2.065(6)  | $Fe-C9$  | 2.068(6)   |
| $Fe-C5$   | 2.024(5)  | $Fe-C10$   | 2.016(6)   |
| $C1 - C11$  | 1,540(7)  | $C11 - C12$  | 1.539(12)  |
| $C6-C12$  | 1.544(7)  | $C1-C2$  | 1.405(8)   |
| $C2-C3$   | 1,408(9)  | $C3-C4$  | 1.431(8)   |
| $C4-C5$   | 1.406(9)  | $C1-C5$  | 1,435(8)   |
| $C1-Fe-C6$<br>$C1-C11-C12$<br>$C3-Fe-C9$<br>$C$ 5-Fe- $C$ 7<br>$C2-C1-C5$<br>$C2-C3-C4$<br>$C2-C1-C11$<br>$C7-C6-C12$ | 88.6(3)<br>105.9(8)<br>120.1(3)<br>101.1(3)<br>108.0(5)<br>107.6(5)<br>130.1(8)<br>136.0(8) | C <sub>11</sub> -C <sub>12</sub> -C <sub>6</sub><br>$C10$ -Fe-C2<br>$C4$ -Fe-C8<br>$C1-C2-C3$<br>$C4-C5-C1$<br>C5-C1-C11<br>C <sub>10</sub> -C <sub>6</sub> -C <sub>12</sub> | 114.5(8)<br>101.1(3)<br>121.2(3)<br>108.6(6)<br>107.4(5)<br>120.8(8)<br>114.3(8) |



Figure 1. Molecular structure of **4a** (vibrational ellipsoids at the 25% probability level).

The most interesting structural feature of **4a** is the tilt of the virtually planar cyclopentadienyl ligands with respect to one another (Figure 1). The tilt angle of  $21.6(4)^\circ$  is comparable to that found for the silicon-bridged [llferrocenophane **1** 



Figure 2. Definition of structural parameters for ferrocenophanes.

*(20.8(5)"),* slightly less than that present in the methylated hydrocarbon-bridged species  $4c (23(1)^\circ)$ . The degree of tilting in 4a can also be appreciated by considering the  $Cp_{\text{centroid}}$ -Fe-Cp<sub>centroid</sub> angle, which is  $164.1(3)^\circ$  compared with  $180^\circ$  in ferrocene,  $163.4(6)^\circ$  for **4c**, and  $164.74(8)^\circ$  in **1**. In **4c**, the displacement of the iron atom from the line joining the two centroids of the cyclopentadienyl ligands is greater than the value in 4a  $(0.432(12)$  vs  $0.225(7)$  Å). By contrast, the angles between the planes of the cyclopcntadienyl ligands and the C(Cp)-C bonds ( $\beta$ ) in **4c** arc 10.8(10)<sup>°</sup> and 8.9(13)<sup>°</sup>, which are smaller than in **4a** (20.1 (3)<sup>o</sup> and 12.7 (3)<sup>o</sup>), and angle  $\delta$  for **4a** (164.1 (3)<sup>o</sup>) shows a considerably greater deviation from 180 $^{\circ}$  than that for **3** ( $\delta$  =  $176.48(3)$ <sup>o</sup>). In **4a**, the displacement of the iron atom from the line joining the two centroids of the cyclopentadienyl ligands (0.225 (7) A) is significantly greater than the value in **3**   $(0.027(3)$  Å). By contrast, the average angles between the planes

of the cyclopentadienyl ligands and the  $C(Cp) - C$ bonds  $(\beta)$  in 4a are 20.1(3)<sup>o</sup> and 12.7(3)<sup>o</sup>, larger than the corresponding  $\beta$  angle in **3**  $(10.8(3)^\circ)$ . Interestingly, the C, bridge in **4a** makes an angle of  $18.4(1)^\circ$  with the plane containing the centroids on the cyclopentadienyl rings and the iron atom and is therefore significantly more twisted than the disilane bridge in **3,** where the corresponding angle is  $8.4(4)^\circ$ .

It is *also* intercsting to compare the structure of the recently reported hydrocarbon-bridged [2]ruthenocenophane  $[Ru(\eta - C_5H_4)_2(CH_2)_2]$  with that of its iron analogue **4a**.<sup>[43, 56]</sup> The larger size that of its iron analogue  $4a^{[43, 56]}$  The larger size of the central ruthenium atom compared with iron results in drastic structural differences. The most interesting difference lies in the tilt anligands in the ruthenium complex (29.6(5)<sup>°</sup>), <br>Figure 3. Solid-state <sup>13</sup>C NMR spectrum of polymer **5a** (\* denotes spinning sidebands). which is ca.  $8.0^\circ$  greater than in the analogous iron species **4a**  $(\alpha = 21.6(4)°)$ . In addition, the

Cp-M-Cp angle  $\delta$  (M = Fe, Ru) for **4a** ( $\delta$  = 164.1(3)<sup>o</sup>) deviates less from 180" than that for the ruthenium analogue  $(\delta = 159.3(2)^{\circ})$ . The displacement of the metal atom in the latter compound from the line joining thc two centroids of the cyclopentadienyl ligands (0.321 *(5)* A) is greater than the corresponding value in  $4a$  (0.225(7) Å).

Synthesis and structural characterization of the poly(ferrocenylethy1ene)s 5a and 5b: Polymerization of **4a** and 4b was achieved by heating these species in the melt at elevated tcmperatures in evacuated, sealed Pyrex tubes. In both cases the tube contents became molten and then rapidly more viscous, and eventually immobile. The polymeric product 5 **b** dissolved in THF and was isolated as a fibrous material by precipitation into methanol. By contrast, the unsubstituted polymer 5 a was found to be essentially insoluble in organic solvents. The colours of 5a and **5b** varied from mustard yellow to yellow-brown. The darker-coloured materials probably contained small amounts of thermal decomposition products (see below).

Although *5* a was insoluble, the polymeric nature of this material was suggested by its film-like appearance on the sides of the tube and by the identification of cyclic oligomers  $[Fe(n-C<sub>5</sub> H_4CH_2$ <sub>2</sub>,  $(x = 2-5)$  in CH<sub>2</sub>Cl<sub>2</sub>-soluble extracts of the material by mass spectrometry. In addition, Soxhlet extraction with hot 'THF over 72 h produced a small amount of soluble material. Gel permeation chromatography (GPC) indicated that 5a possessed a bimodal molecular weight distribution. The first fraction possessed an approximate weight average molecular weight  $(M<sub>n</sub>)$  of  $8.1 \times 10^4$  and a number average molecular weight *(M<sub>n</sub>*) of  $6.6 \times 10^4$  while the second, essentially oligomeric fraction was characterized by values of  $M_w = 4.8 \times 10^3$  and  $M_p = 3.5 \times 10^3$ . Because of the essentially insoluble nature of 5a, this material was characterized by solid-state NMR. The solid-state <sup>13</sup>C NMR spectrum displayed resonances consistent with the formation of a poly(ferrocenylethylene) (Figure 3) with broad resonances at  $\delta \approx 89-91$ ,  $68-72$  and  $35-38$  assigned to C(Cp-*ipso*),  $C(\alpha, \beta Cp)$ , and CH<sub>2</sub>CH<sub>2</sub>, bridging groups respectively. The yields of the **poly(ferrocenylethy1ene)s** 5 a and 5 b were virtually quantitative and no unreacted 4a or **4b** was detected.



The poly(ferrocenylethy1ene) 5 **b,** which was readily soluble in organic solvents, was structurally characterized by multinuclear NMR spectroscopy, UV/vis spectroscopy and elemental analysis. In addition, the molecular weight distribution was analyzed by GPC. The <sup>1</sup>H NMR spectra of 5b (in  $C_6D_6$ ) showed a broad resonance for the cyclopentadienyl protons at  $\delta \approx 3.6-4.1$  and broad resonances assigned to the bridge protons and methyl groups associated with the cyclopentadienyl rings at  $\delta = 2.3$ -2.8 and 1.6-2.1, respectively. The integration ratio of these resonances was 6:4: 6, which was consistent with the assigned structure. The  $^{13}$ C NMR spectrum for 5b was complex, owing to the structural isomerism involving the methyl substituents attached to the cyclopentadienyl rings, but was also consistent with the assigned structure. Significantly, in the <sup>13</sup>C NMR spectrum of **5b**, the resonance associated with the *ipso* carbon of the cyclopentadienyl ring exhibits a slight highfield shift from  $\delta \approx 85.6 - 91.5$  in **4b** to 83.3–88.5 in **5b**, consistent with a structure in which the cyclopentadienyl rings are essentially parallel. The UV/visible data for polymer **5 b** had a low-energy absorption at 440 nm  $(\epsilon_1 = 190 \text{M}^{-1} \text{cm}^{-1})$  very similar to that of ferrocene ( $\lambda_{\text{max}} = 440$  nm;  $\epsilon_1 = 90 \,\text{M}^{-1} \text{cm}^{-1}$ ); this suggests an essentially localized polymer backbone. GPC in THF indicated that **Sb** possessed a bimodal molecular weight distribution. The first fraction had an approximate weight average molecular weight  $(M_{\rm w})$  of  $9.6 \times 10^4$  and a number average molecular weight  $(M_n)$  of  $8.6 \times 10^4$ , while the second, essentially oligomeric fraction was characterized by values of  $M_w = 4.8 \times 10^3$  and  $M_{\rm n} = 3.5 \times 10^3$ .

The existence of a bimodal molecular weight distribution for polymer **5 b** suggests the possibility that multiple mechanisms may operate during the thermal ROP of **4b.** In an attempt to examine the nature of the molecular weight distribution for **5b,**  the polymerization of **4b** was examined more closely. Sealed Pyrex tubes containing **4 b** were heated at 300 *"C* for iutervals of  $0-2.5$  h and the contents of each tube were examined by GPC. Analysis of a tube heated for 30min (at which time the tube contents were viscous but mobile) showed that the polymer present possessed an approximate weight average molecular weight  $(M_{\rm w})$  of  $8.0 \times 10^4$  and a number average molecular weight  $(M_n)$  of  $6.6 \times 10^4$  with no substantial lower molecular weight oligomeric fraction. Analysis of the contents of the tubes which had been heated for 45 min or more (after which time the tube contents were immobile) showed the aforementioned characteristic bimodal molecular weight distribution. In order *to*  determine whether the lower molecular weight fraction arises as a result of thermal decomposition of the polymer, a tube containing a purified sample of polymer 5 **b** was heated for 1 h at 300 'C. Analysis of the tube contents by GPC showed no change in molecular weight or in the molecular weight distribution. Thermogravimetric analysis of polymer **5 b** supports this finding, as this material is stable to weight loss up to ca. 375  $\degree$ C (see below), which is above the ROP temperature. These results provide support for the tentative postulate of two different polymerization mechanisms.

Attempted transition-metal-catalyzed ROP of 4a: Recently, in an effort to induce strained silicon-bridged [l]ferrocenophanes such as 1 to undergo ROP under milder conditions, the use of various transition-metal catalysts was explored. This resulted in the formation of high molecular weight poly(ferrocenylsilane)s (e.g. 2) at room temperature.<sup>[24, 57, 58]</sup> Attempts to extend this chemistry to similarly strained hydrocarbon-bridged [2]metallocenophane systems such as **4a,** however, were, not surprisingly, unsuccessful. Solutions of  $4a$  (in  $C<sub>6</sub>D<sub>6</sub>$ ) were treated with **bis(cyclooctene)rhodium(i)** chloride dimer, and the mixture was stirred under nitrogen with constant monitoring by 'H NMR spectroscopy. Analysis of this mixture after 48 h by both 'H NMR spectroscopy and GPC showed only thc presence of the starting compound **4a** and no signs of oligomeric or polymeric material. Similar results were obtained with  $PtCl<sub>2</sub>$  and platinum divinyltetramethyldisiloxane complex as the catalysts.

**Thermal transition behaviour and morphology of the poly(ferr0 cenylethy1ene)s 5a and 5 b:** In order to obtain information on the conformational flexibility and morphology of the poly(ferrocenylethylene)s, the thermal transition behaviour of **5 a** and **5 b**  was investigated. Whereas differential scanning calorimetry (DSC) showed no evidence for melting transitions for polymer **5b**, a large melting endotherm  $(T_m)$  at 241 °C was detected with the corresponding recrystallization exotherm at 191 *"C* detected on cooling (Figure 4) for polymer Sa. For polymer **Sb,** only a



Figure 4. DSC thermogram for polymer **5a** (scan rate of  $10^{\circ}$ Cmin<sup>-1</sup>).

glass transition with very small change in heat capacity was detected at ca.  $65^{\circ}\text{C}$ , and no apparent  $T_g$  was observed for polymer **Sa.** Comparison of the glass-transition data with data for polyethylene, which has a  $T_{\rm g}$  value of  $-120$  °C, reveals that the inclusion of ferrocene into the polyethylene backbone decreases the skeletal flexibility. Also, comparison of the glass transition temperature for the poly(ferrocenylethy1ene) 5 **b** with that for poly(vinylferrocene)  $(T_g = 184 \degree C)^{59}$  where ferrocene is present as a side group, indicates that the incorporation of a ferrocenyl moiety into the backbone decreases the conformational flexibility less dramatically.

The morphology of the poly(ferrocenylethylene)s was also examined by wide-angle X-ray scattering (WAXS) at 25 *"C.* The WAXS scattering pattern for 5a showed significant order, with a sharp peak corresponding to a d spacing of 5.14 Å (Figure 5). Scattering patterns for **Sb** were broad with no significant signs of long-range order and displayed a broad peak corresponding to a *d* spacing of 6.18 Å superimposed on an amorphous halo (Figure 5, inset).

# **Thermal stability of the poly(ferrocenylethy1ene)s** Sa **and 5 b:**

*u) Tllermogruvimetric anulysis:* Thermogravimetric analysis (TGA) studies of the polymer  $5b$  under  $N_2$  at a heating rate of  $10^{\circ}$ C min<sup>-1</sup> showed that this material was stable to weight loss up to ca. 375°C (Figure 6). Above this temperature two distinct weight-loss processes were detected. An initial weight loss of approximately 48% occurred between 375 °C and 450 °C, with a subsequent weight loss of approximately 20% observed between 450 °C and 500 °C. Further thermolysis up to  $1000$  °C led to very little change in mass. The final char yield for this polymer was 32%. Analogous thermolysis studies on polymer Sa showed similar behaviour with a higher final char yield of 50 %.



Figure 5. Wide-angle X-ray scattering pattern for polymers 5a and 5b (inset) at 25 °C.



Figure 6. Thermogravimetric analysis trace for polymer **5b** obtained at 10 °C min<sup>-1</sup> under N<sub>2</sub>

*h)* Pyrolysis studies in a tube furnace: In order to investigate the thermolysis of poly(ferrocenylethy1ene)s in more detail, studies in a tube furnace were carried out. Polymer Sb was chosen for this work because it could be purified more thoroughly than **5 a,** since the latter material is essentially insoluble in organic solvents. Pyrolysis of *S* **b** was carried out under N, with a 1 h ramp from room temperature to  $600^{\circ}$ C and subsequent thermolysis for 4 h at this temperature. Ceramics were obtained as black, lustrous materials in yields of ca. 30%. Concurrent with the onset temperature for weight loss by TGA *(375"C),* the formation of a red-orange oil was observed on the cooler section of the quartz tube downstream from the polymer sample.

c) *Churucterizution qf the pyrolj.sis products:* The solid pyrolysis product **6b,** formed during the thermolysis of **Sb** at *600°C.*  was readily attracted to a bar magnet and was further investigated by X-ray photoelectron spectroscopy (XPS), high-resolution scanning electron microscopy (SEM) with energy dispersivc Xray microanalysis (EDX) and backscattered electron imaging

Analysis of the ceramic **6b** by XPS indicated the presence of a surface primarily containing carbon (ca.  $87\%$ ) with some oxygen (ca.  $11\%$ ). The presence of oxygen was also noted for the ceramics derived from the pyrolysis of poly(ferrocenylsilane)s and may arise from workup of the pyrolysis products in air.<sup> $[22.29]$ </sup> Analysis of the bulk of the sample by SEM-EDX, with samples of polymer **Sb** for compositional comparisons, showed an iron :carbon :oxygen ratio of approximately 72:24:4. Interestingly, the material was not homogeneous and small regions of higher oxygen concentration (iron: carbon: oxygen ratio =  $70:5:25$ ) were also present (see Figure 7).



Figure 7. a) Scanning electron micrograph of ceramic  $6b$  at  $\times 100$  magnification. b) Back-scattered electron image of ceramic  $6b$  at  $\times 100$  magnification.

The magnetic propcrties of the ceramic **6 b** werc studied by Mössbauer spectroscopy and magnetization measurements. The <sup>57</sup>Fe Mössbauer spectrum of **6b** confirmed the presence of magnetic iron sites with *a* characteristic six-line spectrum arising from the lifting of the degeneracy of the  $I = \pm \frac{1}{2}$  and  $\pm \frac{3}{2}$ states. **A** study of the magnetization of **6b** as a function of applied field gave a hysteresis curve characteristic of a soft ferromagnetic material. **A** study of **6b** by powder XRD indicated the presence of significant amounts of  $\alpha$ -Fe crystallites (sharp peaks at *d* spacings of *2.02(5),* 1.43(3) and 1.17(3) A) together with several broadened peaks of low intensity (Figure 8). Simi-

# **2-Theta** - **Scale**



lar behaviour has been noted for the ceramics derived from the pyrolysis of poly(ferrocenylsilanc)s, where much higher ceramic yields have been achieved.<sup>[21,22,29]</sup>

The liquid produced during the pyrolysis of **5b** was collected in dichloromethane solvent and was analyzed by mass spectrometry. The peaks wcre tentatively assigned to cyclic trimer  $[Fe(\eta - C_5H_3(CH_3)CH_2),]$ , (9) and the linear methylated compounds  $[Fe{\{\eta - C_5H_3Me_2\}\{\eta - C_5H_3(Me)\}(CH_2)]_2}$  (10). Cyclic depolymerization products have been previously detected when poly(ferrocenyldimethylsi1ane) **(2)** is heated at elevated temperatures.<sup>{22,29}</sup>



**Cyclic voltarnrnetry studies of poly(ferrocenylethy1ene) 5 b:** Prcvious cyclic voltammetric studies focussing on the electrochemical behaviour of poly(ferrocenylsi1ane)s such as **2** have shown that there are electronic interactions between the metal atoms of the main chain in these polymers. Thus, poly(ferrocenyldimethylsilanc) **2** exhibits two reversible oxidation waves with a redox coupling  $\Delta E_{1/2}$  of 0.25 V.<sup>[11,25]</sup> To determine whether similar interactions would be present with other elements instead of silicon, cyclic voltammograms of the poly(ferrocenylethylene) **5b** in CH<sub>2</sub>Cl<sub>2</sub> solutions were recorded at a variety of scan rates (Figure 9). Two reversible oxidation waves were observed as *a*  result of slight electronic communication between metal centres along the polymer chain. Initial oxidation of alternating iron

sites at a potential of - *0.25* V was found to increase the oxidation potential of *a* neighbouring iron site to  $-0.16$  V. The presence of a more insulating hydrocarbon bridge dampens clectronic interaction between metal centres along the polymer backbone in polymer **5 b,** which has a smaller redox coupling  $(\Delta E_{1/2})$  of 0.09 V. For 5b. plots of current vs. square root of scan rate were linear over the rangc of scan rates employed *(50-*   $2000 \,\mathrm{mVs^{-1}}$ ) for both oxidation waves; this indicated that the electron transfer was essentially diffusion-controlled (see inset in Figure 9). Electrochemical studies of hy-

drocarbon-bridged biferrocene systems, such as the ethanebridged dimer  $Fc-(CH_2)_2-Fc$ , have shown contrasting behaviour; this compound exhibits only one reversible oxidation wave at  $E_{1/2} = 0.37$  V vs. SCE.<sup>[60]</sup>



Figure 9. Cyclic voltammograms of polymer 5b in CH<sub>2</sub>Cl<sub>2</sub> obtained at scan rate of *a*) 50 and b) 250 mVs<sup>-1</sup> at 25 °C referenced to the ferrocene/ferrocenium couple at  $E = 0.0$  mV. Also shown are plots of  $i_{\rho k}$  vs. the square root of the scan rate for the oxidation **neaks** observed for **Sh.** 

**Mossbauer spectroscopic studies of poly(ferrocenyletby1ene) 5 b:**  A room-temperature Mossbauer spectrum of an air-oxidized sample of Sb was obtained in order to probe the nature of the possible different iron sites in the resulting polymer (Figure 10). The Mössbauer spectrum for the oxidized poly(ferrocenylethylene) S **b** displayed a pair of doublets, with the outer doublet



Figure 10. Mössbauer spectrum of partially oxidized 5b at 25 °C.

having an isomer shift ( $\delta$ ) of 0.382 mm s<sup>-1</sup> and quadrupole splitting value  $(\Delta E_0)$  of 2.33 mms<sup>-1</sup>, while the inner doublet displayed a  $\delta =$  of 0.31 mm s<sup>-1</sup> and  $\Delta E_q$  of 0.69 mm s<sup>-1</sup>, characteristic of Fe<sup>II</sup> and Fe<sup>III</sup> sites, respectively. This implies that the oxidized **5b** has a localized electronic structure on the Mössbauer timescale  $(10^{-7} s)$ .

**Characterization of the oxidation products from the reaction of the poly(ferrocenylethy1ene) 5 b with TCNE:** The preparation of magnetic materials derived from polymer **5b** was somewhat complicated by the presence of very small but variable quantities of magnetic impurities in the polymer. Thesc impurities are presumably pyrolysis products formed during the high temperatures (ca. 300°C) required for the synthesis of 5b.

The unoxidized polymer (even the purest material obtained) showed *a* characteristic, very unsymmetrical ESR signal at an apparent *g* value of ca. 2 at room temperature, but gave no discernible ESR signal below ca. 100 K. Magnetic measurements of samples with greater impurity levels indicate a drop in susceptibility below ca.  $130 \text{ K}$ ; the magnetic susceptibility at low temperature is very small. These results are consistent with the presence of non-ferrocenejferrocenium-derived iron species that undergo a spin crossover at ca. 130 K. The proportion of this impurity is very small, as it does not affect the elemental analysis of the neutral polymer; presumably the species responsible has a rather large moment.

Addition of a dichloromethane solution of one equivalent of TCNE to a dichloromethane solution of **5b** resulted in the formation of two products: a black insoluble material **(11,** ca. 40°/,) was collccted by filtration, and a green precipitate **(12,** ca. 60%,) was obtained by addition of diethyl ether to the filtered dichloromethane solution. Elemental analyses were consistent with stoichiometries close to **5b** [TCNE]<sub>n</sub> for each product.

The only multiple-bond stretching bands observed in the IR spectrum of a nujol mull of 11 were broad intense features centred at 2153 and 2198 cm<sup> $-1$ </sup>; these are indicative of reduced TCNE but at somewhat higher frequency than the isolated TCNE<sup>-</sup> anion.<sup>[61-63]</sup> A plausible explanation is that the materi*al* includes stacks of planar TCNE molecules, which are partially reduced. Compound **11** was also investigated by ESR spectroscopy and SQUID magnetometry. Unfortunately, the magnetic impurity present in the unoxidized polymer appears to end up in this reaction product. Thus, the room-temperature ESR spectrum revealed the characteristic spectrum of the unoxidized polymer, upon which is superimposed a sharp isotropic resonance with  $g = 2.00$ , presumably arising from the cyanocarbon anions. At low temperatures the isotropic resonance completely dominated any signal from the impurity in the polymer. In addition, an axially symmetric ferrocenium spectrum was observed characterized by  $g_{\perp} = 1.70$  and  $g_{\parallel} = 3.76$ . These values are typical for substituted ferrocenium species.<sup>[64-66]</sup> This spectrum is surprisingly sharp given that the sample was a solid and that ferrocenium ions with slightly different substitution patterns and, therefore, with slightly different *g* values, are expected to be present, owing to the variation in the positions of the methyl groups on the cyclopentadienyl rings. The magnetic susceptibility data could not be fitted to the Curie-Weiss law. As magnetic impurities were shown to be present by ESR, no further analysis of the magnetic data was undertaken.

**A** nujol inull of the soluble material **12** shows IR bands at 2148, 2199 and 2218 cm $^{-1}$ , the middle band being particularly intense. This is similar to the spectrum observed for the TCNE $2$ <sup>-</sup> anion,<sup>[67]</sup> although the bands are spread over a wider range of frequencies in **12** and, on average, are at somewhat higher frequency. A very similar IR spectrum is observed in dichloromethane solution, indicating the oligomeric anions do not dissociate in solution. Room-temperature ESR spectra of **12,** either in the solid state or in dichloromethane solution, showed a very poorly resolved isotropic resonance with a *g*  valuc of ca. 2. The characteristic lineshape of the impurity in the unoxidized polymer was not observed; this was taken to indicate that this product is essentially free of the magnetic impurity. The low intensity of the room-temperature radical signal, when compared with spectra of other salts of paramagnetic cyanocarbon species, suggests few anions are paramagnetic. Low-temperature spectra showed the isotropic radical resonance and a ferrocenium spectrum with *g* values almost identical to those from the ferrocenium spectrum of **11.** Spectra were acquired both in dichloromethane and with solid samples; although the solution/glass spectra were slightly sharper, the spectra were 'otherwise very similar, consistent with the anions being un- 'changed when the material is dissolved. Solid-state magnetic susceptibility measurements were made between *5* and 250 K. The data were fitted to the Curie-Weiss law  $(80-250 \text{ K})$  with a Weiss constant  $\Theta$  of  $-11.6$  K (Figure 11). This value is indicativc of significant antiferromagnetic interactions. The magnetic moment is ca. 1.7  $\mu_B$  per iron atom, although this value depends



Figure 11. Variation of molar magnetic susceptibility per Fe  $(\chi, \bullet)$  and effective magnetic moment  $(\mu_{eff}, \times)$  with temperature *(T)* for 12.

greatly upon thc accuracy of the iron analysis. For a fully oxidized polymer with diamagnetic countcrions, one would expect an effective magnetic moment of 2.25  $\mu_{\rm B}$ , if  $\langle g \rangle = 2.60$  as determined by ESR. Below ca. 70 K the plot of reciprocal susceptibility against temperature is no longer linear, and the derived magnetic moment falls to 1.2  $\mu_B$  (Figure 11). This is also consistent with the presence of substantial antiferromagnetic interactions in the material; however, the data show no transition to an ordered antiferromagnetic state.

The formation of two distinct products of similar composition in this reaction is not especially surprising. The products of reactions involving cyanocarbon anions often give more than one product.<sup>[68-70]</sup> Thus, for example, Ward et al. were able to isolate both green  $TCNQ = 7,7,8,8$ -tetracyano-pquinodimethane) and purple  $TCNQ_2^{2-}$  salts of the  $[Cp*Ru (C_6Me_6]^+$  cation from a single reaction mixture.<sup>[70]</sup> The origin *of* the antiferromagnetic interactions in **12** is unknown. However, it is possible that the packing requirements of the cyanocarbon counterions cause ferrocenium ions to be in close proximity with one another.

### **Summary**

The poly(ferrocenylethylene)s  $[Fe(\eta-C_5H_3RCH_2)_2]_n$  5 a and 5 b  $(a: R = H, b: R = Me)$  have been prepared by the thermal ringopening polymerization of the corresponding strained hydrocarbon-bridged [2]ferrocenophanes  $[Fe(\eta-C_5H_3RCH_2)_2]$ . The molecular weight distribution for polymer **Sb** was found to be bimodal in nature with a high molecular weight fraction with  $M_{\rm w} \approx 10^5$  and a low molecular weight fraction with  $M_{\rm w} \approx 10^3$ . This may be the result of the action of two different polymerization mechanisms. A UV/visible spectrum of polymer 5b was consistent with a localized structure for the polymer backbone. The electrochemical behaviour of the polymer 5b was examined by cyclic voltammetry, which revealed that this polymer undergoes two reversible, closely spaced oxidations in  $CH<sub>2</sub>Cl<sub>2</sub>$  solutions at  $-0.25$  and  $-0.16$  V, with a redox coupling  $\Delta E_{1/2}$  of approximately 0.09 V, indicative of only a small interaction between the iron centres. Studies by TGA indicated that poly(ferroceny1ethylene)s are thermally stable to weight loss to about 375 "C under dinitrogen, and at more elevated temperatures yield ferromagnetic ceramic products.

The reaction of the poly(ferrocenylethylene) **5b** with TCNE gave a mixture of two products, 11 and 12, which differ in their solubilities and the degree of oligomerization of the  $TCNE_x^{y-1}$ counterions. Magnetic characterization of **11** was complicated by the presence *of* small amounts of magnetic impurities present in the neutral polymer. Nevertheless, solid-state magnetic susceptibility measurements show significant antiferromagnetic interactions in 12.

**Acknowledgments:** I. M. thanks the Ontario Ccnter for Matcrials Rescarch (OCMR) and the Petroleum Rescarch Fund (PRF), administercd by the American Chemical Society (ACS), for funding, and the Alfred P. Sloan Foundation for a Research Fcllowship (1994--1998). J. M. N. thanks thc University of Toronto for a Simcoe Fellowship, P. N. thanks the Natural Sciences and Engineering Research Council of Canada (NSERC) for a postdoctoral fellowship, and H. R. is grateful to the Deutscher Akademischer Austauschdienst (DAAD) for an overseas exchange award. D. O'H. would like to acknowledge EPSRC for financial support and a studentship (S. B.), and the Royal Society of Chemistry (RSC) Dalton Division for thc Sir Edward Franklin Fellowship. The authors also thank Prof. R. H. Morris for the use of the electrochemical equipment and Prof. G. Ozin for the use of the Mössbauer spectrometer.

Received: August 8, 1996 [F438]

- [1] a) *Inorganic Materials* (Eds.: D. W. Bruce, D. O'Hare), Wiley. Chichester, 1992; b) *Extended Linear Chain Compounds*, Vols. 1-3 (Ed.: J. S. Miller). Pienum, Ncw York, 1982; c) S. L. Ingham, N. J. Long, Angew. Chem. 1994, *106,* 1847; *Angeiv. Cliem. bit. Ed. Enp/.* **1994,** *33,* 1752; d) *L* Orivl. J. L. Serrano, *Adr. Muter.* **1995,** *7,* 348; c) N. *G.* Connelly, W. E. Geiger, *Ah. O~~U~O~JW/. C/iriii.* **1984,** *23.* **1** ; *f,* N. *G.* Connclly. W, E. Geiger. *;hid.* **1985.** *24. 87.*
- (21 D. O'Hare, *Cheni. Soc. Rei~.* **1992,** 121.
- [3] a) J. E. Mark, H. R. Allcock, R. West, *Inorganic Polymers*, Prentice Hall, 1992; b) C. U. Pittman, C. E. Carraher, J. R. Reynolds, in *Encyclopedia of Po/~iiwr Scirncc unit Engineering* (Eds.: H. F. Mark, N. M. Hikales. C. *G.*  Overberger, G. Menges), Wiley, New York, 1989, Vol. 10, p. 541; c) J. E. Sheats, C. E. Carraher, C. U. Pittman, *Metal-Containing Polymer Systems.* Plenum, New York, 1985; d) Inorganic and Metal-Containing Polymeric Mate*rials* (Eds.: J. E. Sheats, C. E. Carraher, C. U. Pittman, M. Zeldin, B. Currell), Plenum, New York. 1989; c) Inorganic and Organometallic Polymers (Eds.: M. Zeldin, K. Wynne. **€1.** R. Allcock), ACS, Washinglon 0. C., **1988,** Vol. 360.
- [4] a) J. S. Miller, **A.** J. Epstein, W. M. ReiFf, *Ace. C/icm. Rc,s.* **1988.** *21.* 114: b) J. S. Miller. A. J. Epstein. W. M. Reiff. *('htvn. Rrv.* **1988. 88,** 201
- *[5]* I. Manners. *Anxciv. C/i(vii.* **1996. iOH.** 1712; *AN~EW. Clieiii ht. Ed Eii~l.* **1996.**  *35,* 1602.
- [6] a) H. M. Nugent, M. Rosenblum, P. Klemarczyk. *J Am. C'hein. Sic.* **1993.** *115.*  3848; b) M. Rosenblum, Adv. Mater. 1994, 6, 159; c) M. Rosenblum, H. M. Nugent, K.3. Jang, M. M. Labes. W. Cahalane, P Klemarczyk. **W** M. Reiff. *Macrnmo/rculc~,s* **1995,28,** 6330; d) R. Arnold. S. **A.** Matchetl, M. Rosenblum. *OJ~g(lnf)i)lL~irl~~i~S* **1988,** 7, 2261 ; e) B. M. Foxman, M. Rosenblum, *ibul.* **1993.**  12, 4805; f) B. M. Foxman, D. A. Gronbech, M. Rosenblum, *J. Organomet. Chem.* **1991,** *413,* 287.
- [7] a) P. F. Bi-andt, T. B. Kauchfuss, *J. Am. Chem. Suc* **1992,** *l/4,* 1926; b) C. P. Galloway, T. B. Rauchfuss, *Angew. Chem.* 1993,  $105$ , 1407; *Angew. Chem. Int. Ed. Engl.* **1993**, 32, 1319; c) D. L. Compton, T. B. Rauchfuss, *Organometallies* 1994, /3, 4367; d) D. L. Compton, P. F. Brandt, T. B. Rauchfuss, D. F. Rosenbaum. *C.* E Zukoski. *Chm. Muter.* **1995,** 7, 2342.
- (81 C. E. Stanton. T. R. Lee. R. H. Grubbs, N. *S.* Lewis, J. K. Pudelski, M. R. Callstrom, M. S. Erickson, M. L. McLaughlin, *Macromolecules* 1995, 28, 8713.
- [9] a) M. E. Wright, E. G. Toplikar, H. S. Lackritz, J. T. Kerney, *Macromolecules* 1994, 27, 3016; b) B. Alonso, M. Morán, C. Casado, F. Lobete, J. Losada, I. Cuadrado, *Chcm Marer.* **1995.** 7. 1440.
- [lo] 0. Nuyken, T. Pohlmann. M. Herberhold. *Mucroniol. Rqi* **1992.** *A20. 11* <sup>1</sup>
- [11] D. A. Foucher, B.-Z. Tang, I. Manners, *J. Am. Chem. Soc.* 1992, 114, 6246.
- **[12]** D. **A.** Foucher, I. Manners, *Mukromol. Chem Rqiid Commuii.* **1993.** *14,* 63.
- [13] C. H. Honeyman, D. A. Foucher, F. Y. Dahmen, R. Rulkens, A. J. Lough, I. Manners, *0rgunonwtuNic.s* **1995.** */I.* 5503.
- [14] J. K. Pudelski, D. P. Gates, R. Rulkens, A. J. Lough, I. Manners, Angew. *C/IWL* **1995,** *107,* 1633; *Angeii. C/icwi Im Ed. Eiigl.* **1995.** *34.* 1506.
- [15] R. Rulkens, A. J. Lough, I. Manners. Angew. Chem. 1996. 108, 1929; Angew. *C/i~m. In/. Ed. Engl.* **1996.** *35,* 1805.
- 1161 For a recent Highlights article on strained metallocenophanes. see M. Hcrberhold, *Angw. Clirm.* **1995,** *107,* 1985; Angeiv. *Chfm liit. Ed. B?g/.* **1995,** *34.*  18.37.
- [I71 D. **A.** Fouchcr, R. Ziembinski, B.-Z. Tang, P. M. Macdonald. **J.** Masaey. *C.* R. Jaeger. **G.** J. Vancso, I. Manners, *Mucroinolccules* **1993,** *26,* 287X.
- [18] D. A. Foucher, C. H. Honeyman, J. M. Nelson, B.-Z. Tang, *I. Manners*, *Angeiv. Clirm.* **1993,** *105,* 1843; *Angew Chein. In/. Ed Di:/.* **1993.** *32.* 1709.
- [I91 D. **A.** Foucher. R. Ziembinski, R. Petersen, J. Pudelski. M. Edwards, Y Ni. J. Massey, C. R. Jaeger, *G. J. Vancso, I. Manners, Macromolecules* 1994, 27,
- 3992.
- [20] J. K. Pudelski, I. Manners, *J. Am. Chem. Soc.* **1995**, 117, 7265.
- **[21]** J. K. Pudelski. R. Rulkcns, D. A. Foucher, **A.** J. Lough. P. M. Macdonald. I. Manners, *Mn~~ronroleculs* **1995.** *28,* 7301
- [22] R. Petersen, D. A. Foucher, B.-Z. Tang, A. J. Lough, N. P. Raju, J. E. Greedan, **1. Manners, Chem. Mater. <b>1995**, 7, 2045.
- [23] J. Rasburn, R. Petersen, T. Jahr, R. Rulkens, I. Manners, G. J. Vancso, Chem. *Muter.* **1995.** 7, 871
- [24] D. L. Zcchel. K. *C.* Hultzsch, R. Rulkens, D. Rslaishis, Y. Ni. J. K. Pudclski, **A.** 1. Lough, 1. Manners. *Orgunomrtu1lic.s* **1996,** *IS,* 1972.
- 1251 I. Manners, *Ah. Orgniioiiir,t. Ciwm.* **1995,** 37, 131.
- [26] a) M. T. Nguyen, A. F. Diaz, V. V. Dement'ev. K. H. Pannell. Chem. Mater. **1993,** *5,* 1389; b) K. H. Pannell. V. V. Dement'ev, 14. LI, F. Cervantes-Lee. M. T. Nguyen. **A.** F. Diaz. *Orgunome/ul/ics* **1994.** *13,* 3644.
- (271 M. Hmyene, A. Yasser, M. Escornc, A. Perchcron-Guegan. F. Garnier, *Ah. iMukr.* **1994.** 6, 564.

- [28] M. J. Drewitt, S. Barlow, D. O'Hare, J. M. Nelson, P. Nguyen, I. Manners, *C%c,nt. Cbnimun.* **1996,** 2153.
- 1291 U.-Z. 'rang, R. Peterscn, D A. Foucher, A. J. Lough, N. Coombs, R. Sodhi. **1.**  Manners, *.L Chcin.* Soc. *hein. in. <'il/iiniiii!.* **1993,** *523.*
- [30] S Barlow. A. L. Roltl. D. O'flare, *Chiwi. Conrniir;/.* **1996.** 257.
- [31] S. Barlow, A. L. Rohl, S. Shi, C. M. Freeman, D. O'Hare, *J. Am. Chem. Soc.* **1996.** *118.* 1578.
- *[31-]* H. Stoeckli-Evan?, **A.** *G.* Oshoine. R. H. Whitcley. *Hizh..* (7ilrinr *Actn* **1976.** .W. 2402
- 133] H. Stoeckli-Evans, A. G. Osborne, R. H. Whiteley, *J. Organomet*, Chem. 1980, *194.* 91.
- **[34]** D Seyferth. H. P. Withers, *O~~i/noi7~~~allicr* **1982,** 1. 1275.
- [35] I. R. Butler, W. R. Cullen, F. W. B. Einstein, S. J. Rettig, A. J. Willis, *Or~onomeru/lic,s* **1983,** 2. 128
- [36] M. Edwards, D. A. Foucher, A. J. Lough, I. Manners, Z. Kristallogr. 1995, *2/11.* X65.
- [37] J. K. Pudelski, D. **A.** Foucher. C. **11.** Honeqman, **A.** J Lough. 1. Manners, <sup>S</sup> Barlow, D. O'Hare, Organometallics 1995, 14, 2470.
- [?XI W kinckh. B.-%. Tang. D. **A.** Voucher. D. B. Zamblc. R. Zicmbinski. **A.**  Lough, I. Manners. *Organometallics* 1993, 12, 823.
- [39] After the publication of ref. [38], Pannell and co-workers also reported the structure of 3: V. V. Dement'ev. F. Cervantes-Lee, L. Parkanyi, H. Sharma, K. H. Pannell, M. T. Nguyen, A. Diaz, Organometallics 1993, 12, 1983.
- 140] M. Burke Laing, K. N. Trueblood, Acta. Crystallogr. 1965, 19, 373.
- [41] J. M. Nelson, H. Rengel, I. Manners, *J. Am. Chem. Soc.* 1993, 115, 7035.
- 1421 Alternative nanies for polymers such as **5a** ;ind **5b** include poly(ferrocenylcne- ethylene)s, poly(ferrocenyleneethane)s and poly(ferrocenediylethane)s. A similar nomenclature situation arises for polymers such as 2.
- 143] J. M. Nelson, A. J. Lough, I. Manners, Angew. Chem. 1995, 106, 1019; Angew. *<'/n~iii Tn/. GI. Engl.* **1994,** *33,* 9x9.
- [44] D. A. Foucher, R. Ziembinski, R. Rulkens, J. M. Nelson, I. Manners, in *Inorfiunk iitidOrfiiiiiCin~i~f~iIl~(. Pdtwvrc lI(Edy* ' P. Wision-Neilwn, Ii. K. Allcock. K J. Wynne). ACS. Washington D. C.. **1994,** Vol. 572. p. 442.
- [45] J K. Pudelskt. D. **A.** Foucher. C. H. Honeyman, P. M. Macdonald. I. Manners, S. Barlow, D. O'Hare, *Macromolecules* 1996, 29, 1894.
- 146) C. **S.** Cundy, C. Eaborn, M. F. Lappcrt, *J.* **Oi:ycniimc/.** *Chrni.* **1972.** *44.* 291
- [47] S. Collins, Y. Hong, N. J. Taylor, Organometallics 1990, 9, 2695.
- [48] E. W. Neuse, H. Rosenberg, *J. Macromol. Sci. Rev. Macromol. Chem.* 1970, C4, 1.
- [49] E W. Neuse, E. Quo, *J.* Po/~ni. *Sci.* **1965. ,43.** 1499.
- *[50]* A. Lutttnghaus, W. Kullick, Angiw *C'hox* **1958,** *70. 43X.*
- [Sl] **tl.** *L.* Lentzner, W *E.* Watts. *('hen Cm77mim.* **1970,** 906.
- 1521 H. I,. Lentzner. W. E. Watts. *Tetraheilron* **1971.** 27, 4343.
- [53] **A.** *G.* Osbornc. **K.** H. Whilcley, R. E. Meads, *.I* Or~inwwr. *('ho71* **1980.** *193.*  -345.
- [54] R. B. Abramovitch, J. J. Atwood, M. L. Good, B. A. Lampert. *Inorg. Chem.* **1975.** */4.* 3085.
- 1581 K. I-Iafner. C. Mink. H.J. Lindner, *Angw. Chcm. In/. Ed. EnxI.* **1994,** *33.*  1479.
- 1561 **13erbcrhold** ct al. have recenlly reported the synthesis and structure of  $[Ru(\eta-C_5H_4CMe_2)_2]$  (average tilt angle  $(\alpha) \approx 31^\circ$ ), see M. Herberhold, T. Bärtl, %. *Nii/iirfor,\c/i.* **1995,** *50h,* 1692.
- **[57]** Y Ni. R. Rulkens, J. K. Pudelaki. 1. Manners, *Mii/unnm/.* ('hiwi *RirpirI* C'oni-1i11111. **1995.** *16,* 637.
- [58] N. P. Reddy, H. Yamashita, M. Tanaka, *J. Chem. Soc. Chem. Commun*. 1995, *7263.*
- [59] C. U. Pittman, J. C. Lai, D. P. Vanderpool, M. Good, R. Prado, Macro-*/i7lJ/c'('ll/<,.\* **1970,** *6, 146.*
- [60] W. H. Morrison, S. Krogsrud, D. N. Hendrickson, *Inorg. Chem.* **1973**, 12, 1998.
- <sup>1611</sup>0. W. Webster. W Mahler. K. *E.* Uenson. *J Ani. C/wn/. So?* **1962.** *84.* 3678.
- *[62] J. Stanley, D. Smith, B. Latimer, J. P. Devlin, J. Phys. Chem. 1966, <i>70, 2011*.
- 1631 *M* F. Kcitig, R. M **Wiiig,** in or^?. *C:hn.* **1969.** *X.* 2685.
- [64] D. M. Duggan, D. N. Hendrickson, *Inorg. Chem.* 1975, 14, 955.
- *[h5]* R. Piins. **A.** R Korswagen, *J. Org~inomrr Chen7.* **1970,** *3,* C74.
- [66] J. S. Miller, D. T. Glatzhofer, D. M. O'Hare, W. M. Reiff, A. Chakraborty, **A.** J. Epstetn. *Inorfi. Chfw7.* **1989.** *28.* 2930.
- [67] J. S. Miller, D. M. O'Hare, A. Chakraborty, A. J. Epstein, *J. Am. Chem. Soc.* **1989.** *I/ I,* 7853.
- [68] J. S. Miller, J. H. Zhang, W. M. Reiff, D. A. Dixon, L. D. Preston, A. H. Reis. E. Gebert. M. Extine, J. Troup. A. J. Epstein, M. D. Ward, *J. Phys. Chem.* 1987, *91.* 4344
- 109] M. D. Ward, D. C. Johnson, *Inorg. Chem.* **1987**, 26, 4213.
- *[70] M. D. Ward, P. J. Fagan, J. C. Calabrese, D. C. Johnson, J. Am. Chem. Soc.* **1989.** *//1.* 1719.
- Data were collected on an Enraf Nonius CAD-4 diffractomoter with graphitemonochromated Mo<sub>Ke</sub> radiation ( $\lambda = 0.71073$  Å). The intensities of three standard reflections measured every 120 minutes showed no decay. The data were corrected for Lorentz and polarization effects and an empirical absorption correction was applied (G. M. Sheldrick, SHELXA-90, Program for Absorption Correction, University or Gottingen (Germany), **1990).** Minimum and maximum absorption corrections were 0.2547 and 0.8874. The structures were solved and refined with the SHELXTIAPC package (G. M. Sheldrick. SHELXTL\PC, Siemens Analytical X-ray Instruments Inc., Madison, Wisconsin **(USA)**, 1994). Refinement was by full-matrix least-squares on  $F^2$  using all data (negative intensities included). The weighting scheme was  $w =$  $1/[\sigma^2(F_\text{o}^2) + (0.0549 P)^2 + 7.08 P]$  where  $P = (F_\text{o}^2 + 2F_\text{o}^2)/3$ . Hydrogen atoms included in calculated postions. The -CH<sub>2</sub>-CH<sub>2</sub>- bridge is disordered over two sites with relative occupancies C11A/C12A:C11B/C12B of 60:40.